

(19)



Europäisches Patentamt

European Patent Office

Office européen des brevets



(11)

EP 0 933 826 A1

(12)

EUROPEAN PATENT APPLICATION

(43) Date of publication:
04.08.1999 Bulletin 1999/31

(51) Int. Cl. 6: H01M 8/02

(21) Application number: 99102024.9

(22) Date of filing: 01.02.1999.

(84) Designated Contracting States:
AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU
MC NL PT SE
Designated Extension States:
AL LT LV MK RO SI

(30) Priority: 03.02.1998 JP 2180198

(71) Applicant:
MATSUSHITA ELECTRIC INDUSTRIAL CO., LTD.
Kadoma-shi Osaka (JP)

(72) Inventors:
• Uchida, Makoto
Hirakata-shi (JP)

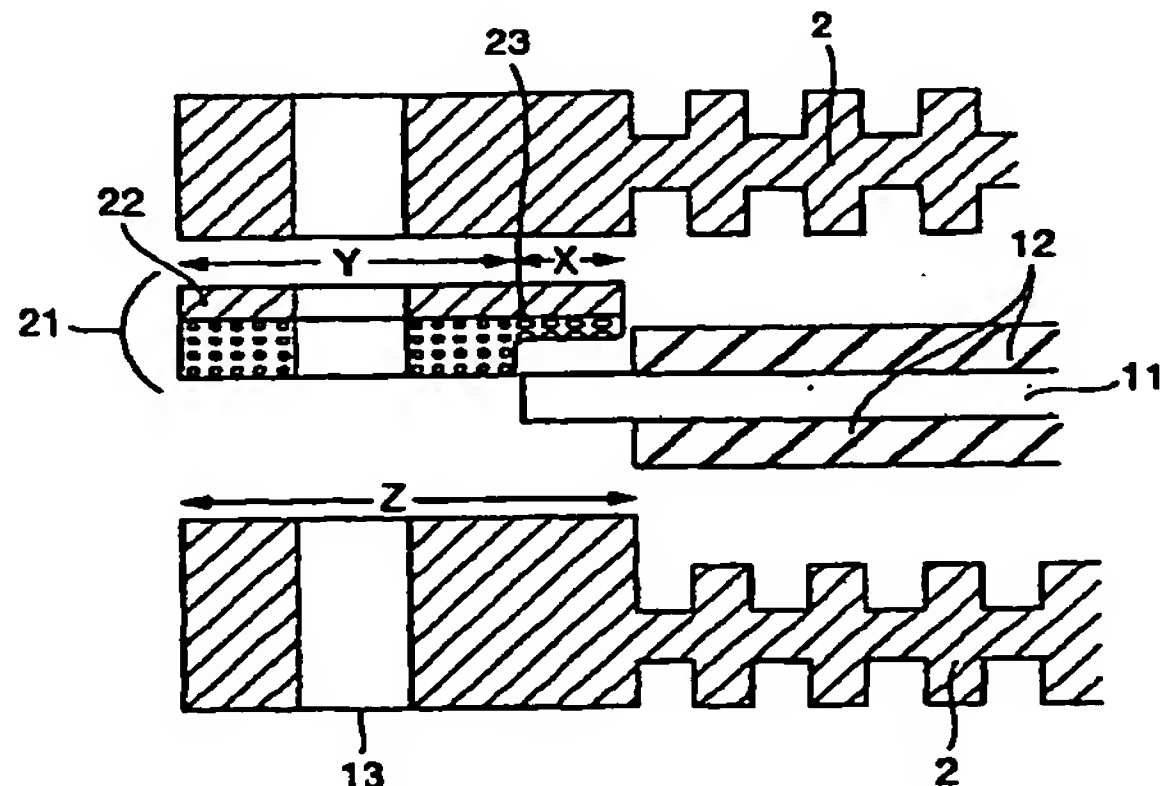
- Fukuoka, Yuko
Fushimi-ku, Kyoto-shi (JP)
- Sugawara, Yasushi
Neyagawa-shi (JP)
- Ohara, Hideo
Kadoma-shi (JP)
- Eda, Nobuo
Hirakata-shi (JP)

(74) Representative:
Patentanwälte
Leinweber & Zimmermann
Rosental 7,
II Aufgang
80331 München (DE)

(54) Polymer electrolyte membrane fuel cell and seal assembly therefor

(57) A polymer electrolyte fuel cell which uses a gasket which comprises an elastomer layer that is inexpensive, highly resistant to chemicals, particularly to acids, and exhibits a high sealability and an adhesive layer provided to the elastomer layer and which gasket is easy to position and easy to assemble. The fuel cell comprises unit cells each comprising a positive electrode (12), an electrolyte plate (11) and a negative electrode (12) and gaskets (21) each arranged at the circumferential part of the unit cell alternately stacked via a separator (2) placed therebetween, wherein the gasket comprises an elastomer layer (23) and an adhesive layer (22), said elastomer layer being adhered to at least one side of the separator via said adhesive layer.

FIG.1A



EP 0 933 826 A1

Description**BACKGROUND OF THE INVENTION**5 **1. Field of the Invention**

[0001] The present invention relates to a fuel cell which uses as a fuel such a reducing agent as pure hydrogen or reform hydrogen obtained from methanol or a fossil fuel and uses air, oxygen or the like as an oxidizing agent. In more particular, it relates to a gasket used for a polymer electrolyte fuel cell.

10

2. Description of Related Art

[0002] It is known that in a polymer electrolyte fuel cell, in cases where for example the cell uses a cation exchange membrane, which is a proton conductor, as the polymer electrolyte and hydrogen and oxygen are introduced thereinto respectively as the fuel and the oxidizing agent, reactions represented by the following formulas (1) and (2) take place.



20

[0003] In the negative electrode, hydrogen dissociates into protons and electrons. The proton moves through the cation exchange membrane toward the positive electrode. The electron moves through electroconductive separator plates, cells stacked therewith in series and further an external circuit and reaches the positive electrode, whereby electricity is generated. In the positive electrode, on the other hand, proton which have moved and reached through the cation exchange membrane, electrons which have moved and reached through the external circuit and oxygen introduced from outside react with one another to form water. Since the reaction is accompanied by heat generation, electricity, water and heat are generated from hydrogen and oxygen, as a whole.

[0004] A polymer electrolyte fuel cell differs greatly from other fuel cells in that its electrolyte is composed of an ion exchange membrane, which is a solid polymer. The ion exchange membrane used include, for example, perfluorocarbon sulfonic acid membrane (Nafion, a trade name, mfd. by Du Pont de Nemours, E.I. Co., USA). In order to show a sufficient proton conductivity, the membrane needs to be in a sufficiently hydrated condition. The hydration of the ion exchange membrane may be effected, as described for example in J. Electrochem. Soc., 135 (1988), p. 2209, by passing the reaction gas through a humidifier to introduce water vapor into the cell and thereby to prevent the drying of the ion exchange membrane. Sealing of the each cell may be effected, as described for example in J. Power Sources, 29 (1990), p. 367, by a method wherein the area of the ion exchange membrane is made larger than the electrode area and the circumferential part of the ion exchange membrane which is not bonded to the electrode is held by the upper and the lower gaskets between them.

[0005] The materials generally used for the gasket include glass fiber fabric coated with polytetrafluoroethylene (Teflon, a trade name, mfd. by Du Pont de Nemours, E.I. Co., USA) and fluororubber. USP No. 4,826,741 discloses the use of silicone rubber and fluororubber.

[0006] Fig. 2 shows an exterior view of a common stack-type polymer electrolyte fuel cell. Separator plates 2 formed of a conductive material, such as glassy carbon, and internal cells (not shown in the Figure) whose circumferential parts are held between insulating gaskets 1 are stacked alternately. A copper-made current collecting plate 3 is closely stuck to the outermost separator plate to form a stack as a whole. The stack is put between stainless steel end plates 5 via insulating plates 4 and the two end plates are bound fast with bolts and nuts. In the Figure, numeral 6 indicates a hydrogen inlet, 7 a hydrogen outlet, 8 an oxygen inlet, 9 an oxygen outlet and 10 a water discharge drain.

[0007] Fig. 3 shows a sectional view of an internal cell of a common stack-type cell. Electrodes 12 are bonded to the both sides of an ion exchange membrane 11 of the center to form an assembly. Grooved separator plates 2 are positioned at the upper and lower sides of the assembly. The ion exchange membrane 11 has a larger area than the electrode 12, and the circumferential part of the membrane is held by gaskets 1 between them to seal each cell and insulate the separator plates from each other. When, as shown in the Figure, a gas path 13 is provided inside the stack according to necessity (that is, in the case of internal manifold type), the gasket serves also to seal the gas path. The separator plate 2 provided with grooves may have various structures; for example, a porous grooved plate is fixed into the groove, or a wire mesh is used in the groove.

55

BRIEF SUMMARY OF THE INVENTION

[0008] However, the above-mentioned prior methods have various problems. When the respective cells are stacked,

in the operation of placing the gasket accurately on the separator plate and holding the assembly of the ion exchange membrane 11 and the electrode 12 by the gaskets between them, the gasket, which is soft and in the form of sheet, can be difficultly positioned and hence gives a poor operation efficiency, or it is apt to give rise to defective seal due to mis-positioning.

5 [0009] Further, when a high pressure gas is used, the gasket tends to get away to the outside of the stack.
 [0010] To solve the above-mentioned problems, the gasket used in the present invention is given a structure comprising an elastomer layer which is inexpensive and highly resistant to chemicals, particularly to acids, and exhibits a high sealability and an adhesive layer. By virtue of the structure, a polymer electrolyte fuel cell having a large economical advantage which uses the gasket that is easy to position and easy to assemble is provided.
 10 [0011] Thus, the fuel cell of the present invention is a fuel cell which comprises unit cells each comprising a solid polymer ion exchange membrane and a positive and a negative electrodes formed on the both sides of the membrane and gaskets each arranged at the circumferential part of the unit cell alternately stacked with each other via a separator placed therebetween, wherein the gasket comprises an elastomer layer and an adhesive layer, said elastomer layer being adhered to at least one side of the separator via said adhesive layer. Accordingly, at the time of assembling a cell stack, since the gasket can be adhered to the separator, mispositioning of the gasket is prevented from occurring.
 15

BRIEF DESCRIPTION OF THE SEVERAL VIEWS OF THE DRAWINGS

[0012]

20 Fig. 1A is a sectional view of a cell in one embodiment of the present invention.
 Fig. 1B is a sectional view of a cell in one embodiment of the present invention.
 Fig. 1C is a sectional view of a cell in one embodiment of the present invention.
 Fig. 1D is a sectional view of a cell in one embodiment of the present invention.
 25 Fig. 2 is an exterior view of a polymer electrolyte fuel cell of the prior art.
 Fig. 3 is a sectional view of a prior cell.

DETAILED DESCRIPTION OF THE INVENTION

30 [0013] According to the present invention, there is provided a fuel cell which comprises unit cells each comprising a solid polymer ion exchange membrane and a positive and a negative electrodes formed on the both sides of the membrane and gaskets each arranged at the circumferential part of the unit cell alternately stacked with each other via a separator placed therebetween, wherein the gasket comprises an elastomer layer and an adhesive layer, said elastomer layer being adhered to one side of at least one separator via said adhesive layer.
 35 [0014] The ion exchange membrane has a larger than the positive electrode and the negative electrode. Consequently, the unit cell has an exposed ion exchange membrane part.
 [0015] The gasket has a dimension sufficient to cover at least the exposed part of the ion exchange membrane. It may further has a dimension which allows provision of a gas path.
 [0016] The gasket comprises an elastomer layer and an adhesive layer.
 40 [0017] According to the above-mentioned structure, the gasket can be adhered to the separator at the time of assembling a cell stack, so that mispositioning of the gasket does not occur and the assembling operation can be proceeded speedily. Furthermore, since the elastomer layer is adhered to the separator plate via the adhesive layer, even when a high pressure gas is used, the elastomer layer does not get away to the outside by virtue of the adhesive force between the elastomer layer and the separator plate.
 45 [0018] Since the working temperature of a polymer electrolyte fuel cell is not higher than 150°C, the elastomer used therein may be various elastic materials, including fluororubber. However, since an ion exchange membrane has sulfonic acid groups as its exchange group and hence is acidic and moreover water is formed in the cell and the reaction gas is humidified, the elastomer needs to be resistant to acids, water vapor, hot water, or the like. Any desired materials may be adopted so long as the above-mentioned conditions of being resistant to heat, acid, water vapor, hot water or
 50 the like are satisfied.
 [0019] However, fluororubbers are expensive and silicon rubbers, in some cases, gradually undergo scission of the siloxane linkage due to the acidity of the sulfonic acid group of the ion exchange membrane and resultant degradation. Therefore, elastomers preferred for use are olefinic rubbers and blend rubbers comprising olefinic rubbers. The blending ratio is not particularly restricted and may be selected according to necessity. Olefinic elastomers which contain no or substantially no unsaturation bond in the polymer main chain are excellent in chemical resistance, heat resistance and weather resistance as compared with diene rubbers, which have double bonds in the main chain, such as isoprene rubber, butadiene rubber, nitrile rubber and chloroprene rubber. Olefinic elastomers, as compared with fluororubbers and silicone rubbers, are inexpensive and excellent in weather resistance.

[0020] Olefinic rubbers preferably used include ethylene-propylene rubber, acryl rubber, butyl rubber and halogenated butyl rubber.

[0021] The adhesive used is not particularly restricted but it is preferably acrylic solvent type adhesive, polyisobutylene rubber type adhesive and isobutylene-isoprene rubber type adhesive. The thickness of the elastomer layer and of the adhesive layer need to be sufficient to achieve insulation and sealing between adjacent separators while absorbing the thickness of the ion exchange membrane. The thickness is preferably 10 - 300 μm for the adhesive layer and 100 - 1000 μm for the elastomer layer. The thickness of the part of the gasket which comes in contact with the ion exchange membrane may be reduced as far as the thickness of the ion exchange membrane as the limit. Similarly, the thickness of the part (Z) of the separator which comes in contact with a laminate of the gasket and the unit cell may be changed as shown in Fig. 1A.

Examples

[0022] The fuel cell of the present invention is explained with reference to Drawings.

[0023] Fig. 1(a) is a sectional view of a cell of Example 1 of the present invention. In the Figure, the gasket 21 is a product obtained by adhering an elastomer layer 23 of olefinic ethylene-propylene rubber (EPDM) of 0.7 mm thickness to one side of a separator plate via an adhesive layer 22. The gasket 21 of the present invention can achieve both the sealing between separator plates and the sealing between an ion exchange membrane and a separator while, as shown in Fig. 1(b), absorbing the thickness of the ion exchange membrane 11 by virtue of the part (X) which comes in contact with the ion exchange membrane 11 being compressed to a more extent than the part (Y) which is held between two separator plates 2. In the case of a gasket comprising an elastomer layer alone, which is very soft, when the internal pressure of the cell and the gas path becomes high the gasket shifts to the outside and is blown through. In the case of the gasket of the present invention, on the other hand, the elastomer layer is prevented from shifting by the adhesive force of the adhesive layer and the gasket is not blown through. At the time of assembling a cell stack, moreover, since the gasket can be adhered to the separator plate beforehand, mispositioning of the gasket does not occur at the time of assembling and the operation can be proceeded speedily. The gasket is also excellent in heat resistance and acid resistance and is not affected in the long term performance test of the fuel cell.

[0024] Though a method of sealing the ion exchange membrane from one direction by using one piece of gasket which has one adhesive layer was shown in Fig. 1(a), similar effects can be obtained by using a gasket which has two adhesive layers as shown in Fig. 1(c) or by using two pieces of gaskets and holding the ion exchanging membrane between the gaskets as shown in Fig. 1(d).

Example 1

[0025] A fuel cell was prepared according to the structure of Fig. 1(a). The gasket 21 was one obtained by adhering an elastomer layer 23 of olefinic ethylenepropylene rubber (EPDM) of 0.7 mm thickness to one side of a separator plate via an adhesive layer 22. The gasket of the present invention was prevented from shifting of the elastomer layer by the adhesive force of the adhesive layer and was not blown through. At the time of assembling a cell stack, moreover, since the gasket could be adhered to the separator plate beforehand, mispositioning of the gasket did not occur at the time of assembling and the operation could be proceeded speedily.

[0026] The EPDM used was kept in contact with the ion exchange membrane in a hot water of 80°C for 3 months to examine the hot water resistance and acid resistance of the elastomer material. The EPDM showed no change and hence showed a high durability. It was also not affected in the 5000 hours long-term performance test of the fuel cell.

Example 2

[0027] A fuel cell was prepared according to the same structure as in Example 1 except for changing the elastomer to olefinic butyl rubber (IIR). Similarly to EPDM, the butyl rubber showed no change in the above-mentioned heat resistance and acid resistance tests, and showed no degradation in the long term performance test of the fuel cell.

Example 3

[0028] A fuel cell was prepared according to the same structure as in Example 1 except for changing the elastomer to olefinic acryl rubber (ACM). Similarly to EPDM, the acryl rubber showed no change in the above-mentioned heat resistance and acid resistance tests and showed no degradation in the long term performance test of the fuel cell.

Example 4

5 [0029] A fuel cell was prepared according to the same structure as in Example 1 except for changing the elastomer to olefinic halogenated butyl rubber (X-IIR). Similarly to EPDM, the halogenated butyl rubber showed no change in the above-mentioned heat resistance and acid resistance tests and showed no degradation in the long term performance test of the fuel cell.

Referential Example 1

10 [0030] A fuel cell was prepared according to the same structure as in Example 1 except for changing the elastomer to diene-type nitrile rubber (NBR). In the above-mentioned heat resistance and acid resistance tests, the ion exchange membrane discolored and the rubber was found to have lowered its elasticity.

Referential Example 2

15 [0031] A fuel cell was prepared according to the same structure as in Example 1 except for changing the elastomer to diene-type chloroprene rubber (CR). After 46 days in the above-mentioned heat resistance and acid resistance tests, swelling of 160% or more was observed and the rubber was found to have lowered its elasticity.

20 Referential Example 3

25 [0032] A fuel cell was prepared according to the same structure as in Example 1 except for changing the elastomer to silicone rubber. After 46 days in the above-mentioned heat resistance and acid resistance tests, the part of the rubber which had been in contact with the ion exchange membrane was observed to have degraded and changed into fine powder. In the long term performance test of the fuel cell, the contact part of the silicone gasket with the ion exchange membrane was found to have changed into silica (SiO_3)-like fine powder.

Referential Example 4

30 [0033] A fuel cell was prepared according to the same structure as in Example 1 except for changing the elastomer to butadiene-type styrene-butadiene rubber (SBR). After one month in the above-mentioned heat resistance and acid resistance tests the rubber was found to have swollen and degraded.

[0034] The structural formulas of the elastomers of Examples and Referential Examples are summarized in Table 1.

35

40

45

50

55

Table 1

| | Elastomer | Structural Formula | Random Copolymer |
|-----------------------|---------------------------------------|--|---|
| Example 1 | Ethylene-propylene Rubber EPM EPDM | $\begin{array}{c} \text{CH}_3 \\ \\ \text{---CH}_2\text{---CH}_2\text{---}(\text{CH}_2\text{---CH---})_n\text{---} \\ \\ \text{CH}_3 \end{array}$ $\begin{array}{c} \text{CH}_3 \\ \\ \text{---CH}_2\text{---CH}_2\text{---}(\text{CH}_2\text{---CH---})_n\text{---} \\ \\ \text{---Cyclohexadiene---} \\ \\ \text{---CH---CH}_3 \end{array}$ | $\begin{array}{c} \text{CH}_3 \\ \\ \text{---CH}_2\text{---CH}_2\text{---}(\text{CH}_2\text{---CH---})_n\text{---} \\ \\ \text{CH}_3 \end{array}$ Ditto ENB-Type |
| Example 2 | Butyl Rubber IIR | $\begin{array}{c} \text{CH}_3 \\ \\ \text{---CH}_2\text{---C---}(\text{CH}_2\text{---C=CH---CH}_2)_n\text{---} \\ \\ \text{CH}_3 \end{array}$ | Isoprene Content 2 mol% or less |
| Example 3 | Acryl Rubber ACM ANM | $\begin{array}{c} \text{CH}_2\text{---CH---}(\text{CH}_2\text{---CH---})_n\text{---} \\ \\ \text{O=C---OR} \end{array}$ $\begin{array}{c} \text{CH}_2\text{---CH---}(\text{CH}_2\text{---CH---})_n\text{---} \\ \\ \text{OCH}_2\text{CH}_2\text{Cl} \end{array}$ | R: Ethyl, Butyl, etc. |
| Referential Example 1 | Nitrile Rubber NBR | $\begin{array}{c} \text{CH}_2\text{---CH=CH---CH}_2\text{---}(\text{CH}_2\text{---CH---})_n\text{---} \\ \\ \text{CN} \end{array}$ | Random Copolymer |

| Structural Formula | | |
|-----------------------|---------------------------------|--|
| Referential Example 2 | Elastomer Chloroprene Rubber CR | $ \begin{array}{c} \text{C1} \\ \\ \text{+CH}_2-\text{C=CH-CH}_2-\text{+} \end{array} $ <p>High-trans</p> |
| Referential Example 3 | Silicone Rubber Q | $ \begin{array}{c} \text{CH}_2 \\ \\ \text{+Si-O-+} \quad \text{+Si-O-+} \\ \quad \\ \text{CH}_3 \quad \text{CH}_3 \end{array} $ <p>Methylvinylsilicone Rubber (VMQ)</p> |
| Referential Example 4 | Styrene-Butadiene Rubber SBR | $ \begin{array}{c} \text{CH=CH}_2 \\ \\ \text{+Si-O-+} \quad \text{+Si-O-+} \\ \quad \\ \text{CH}_2\text{CH}_3\text{CF}_3 \quad \text{CH}_3 \end{array} $ <p>Fluorosilicone Rubber</p> <p>  Random Copolymer </p> |

[0035] Similar effects were obtained also when materials obtained by blending the above-mentioned elastomers with each other or blending the elastomer(s) with other elastomers were used.

[0036] As set forth above, according to the present invention, a polymer electrolyte fuel cell having a large economical advantage can be provided which uses a gasket which comprises an elastomer layer that is inexpensive, highly resistant to chemicals, particularly to acids, and exhibits a high sealability and an adhesive layer provided to the elastomer layer and which gasket is easy to position and easy to assemble.

5

Claims

1. A fuel cell which comprises unit cells each comprising a solid polymer ion exchange membrane and a positive and a negative electrodes formed on the both sides of the membrane and gaskets each arranged at the circumferential part of the unit cell alternately stacked with each other via a separator placed therebetween, wherein the gasket comprises an elastomer layer and an adhesive layer, said elastomer layer being adhered to at least one side of the separator via said adhesive layer.
2. The fuel cell according to claim 1 wherein the elastomer consists essentially of olefinic rubber or blend rubber of olefinic rubbers.
3. The fuel cell according to claim 2 wherein the olefinic rubber is at least one member selected from the group consisting of ethylene-propylene rubber, acryl rubber, butyl rubber and halogenated butyl rubber.

20

25

30

35

40

45

50

55

FIG.1A

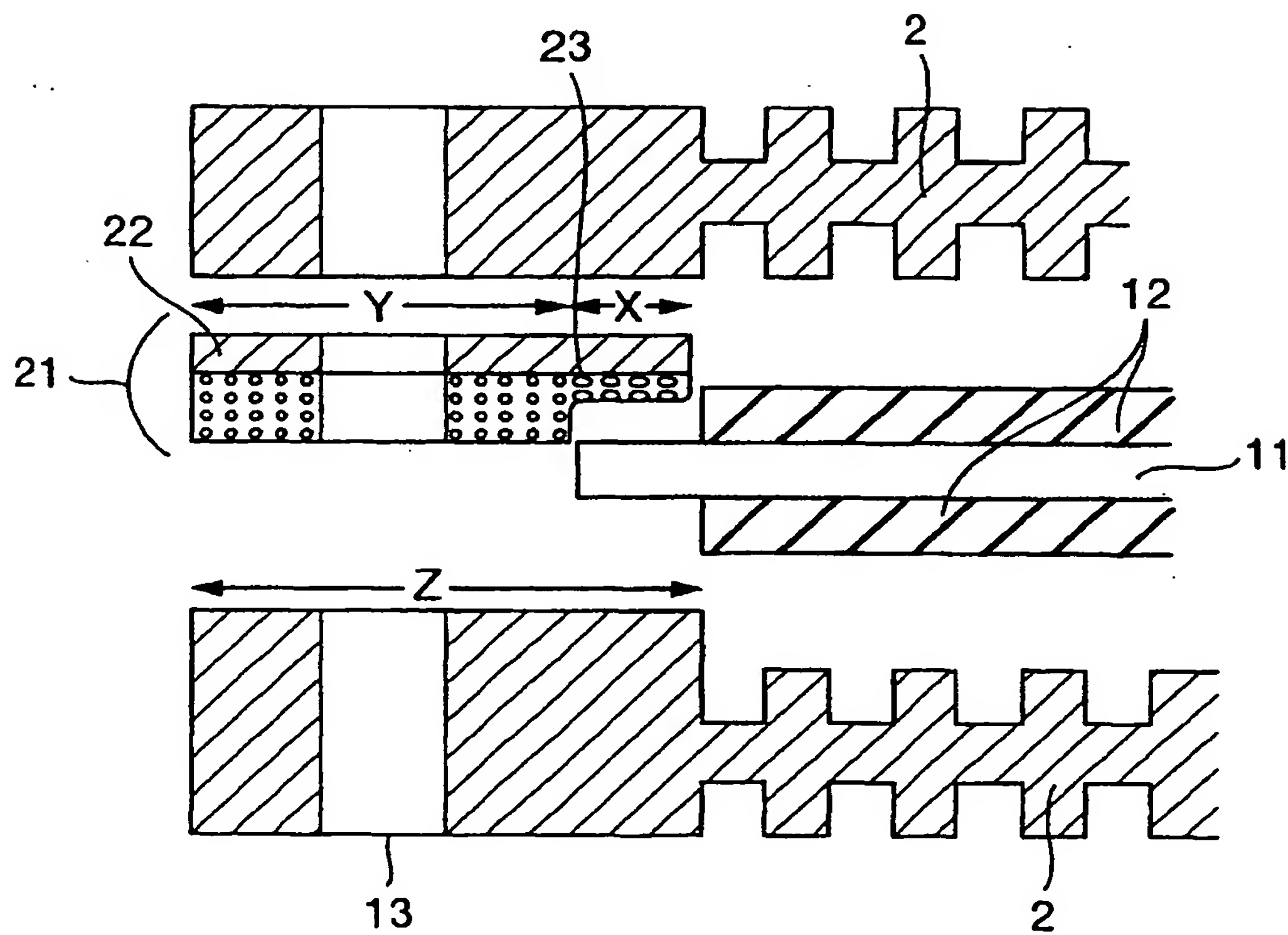


FIG.1B

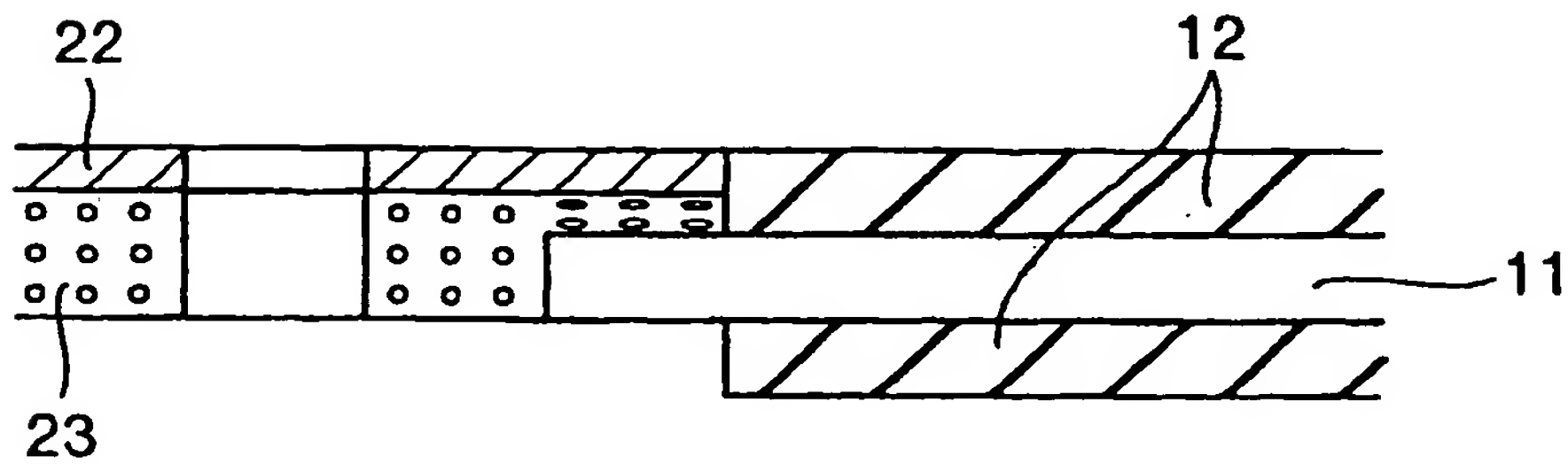


FIG.1C

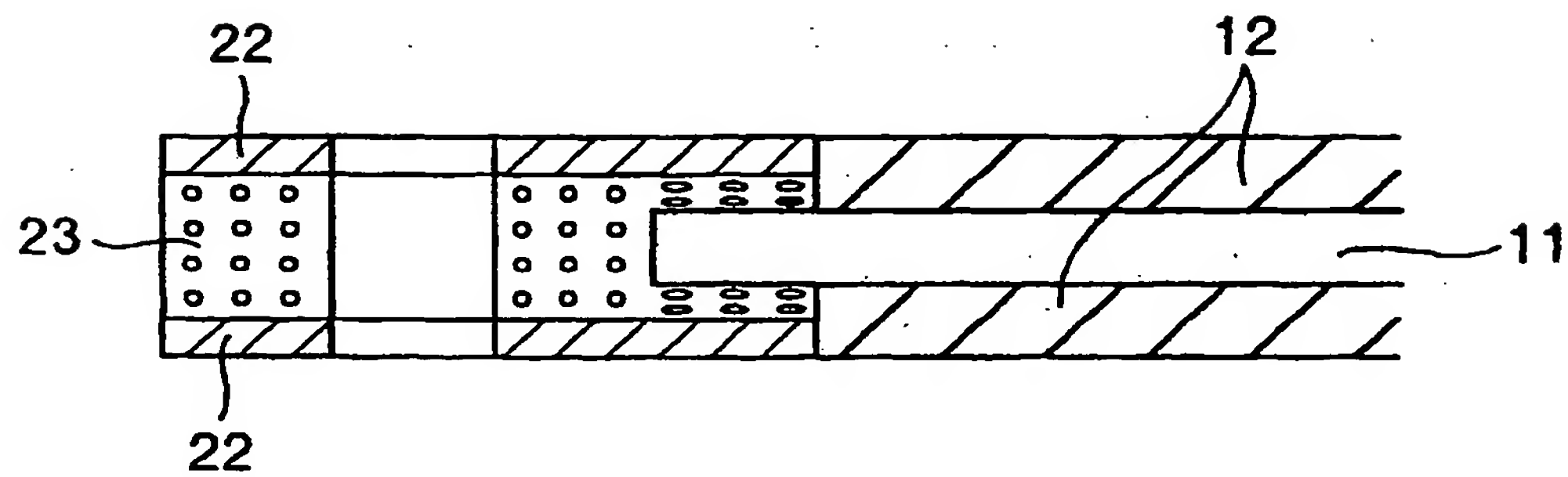


FIG.1D

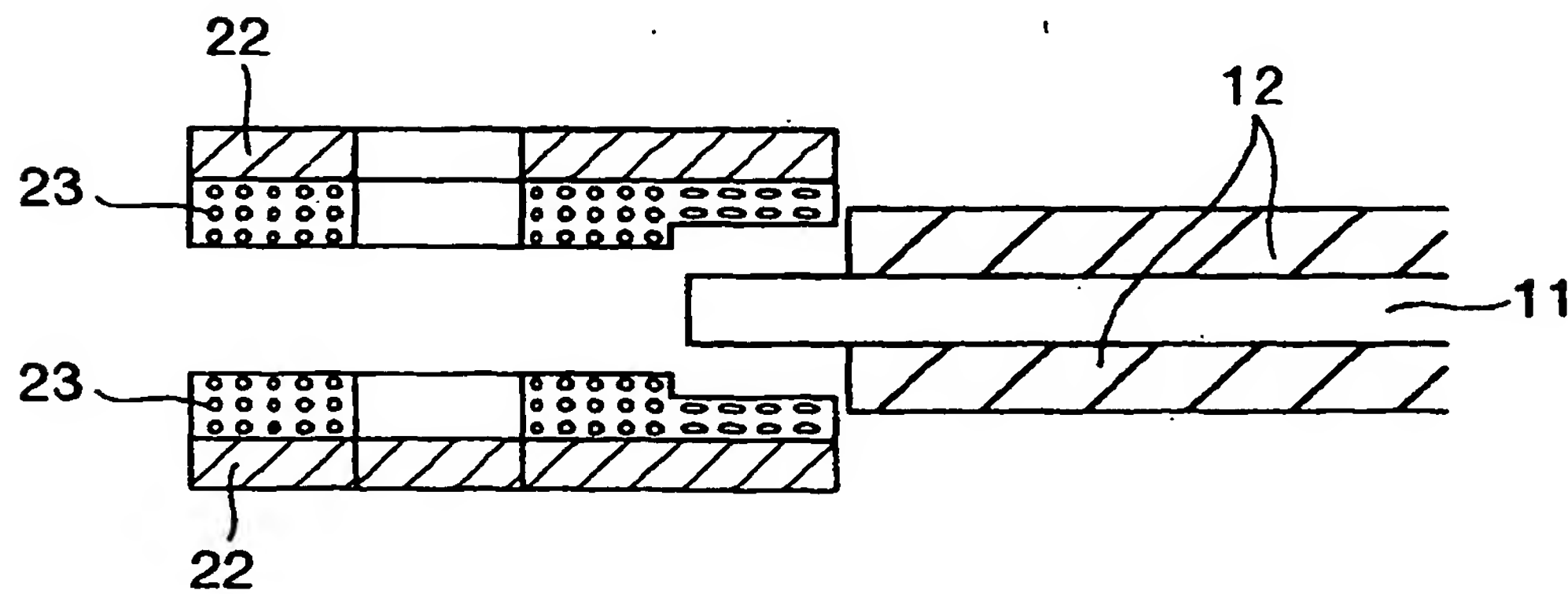
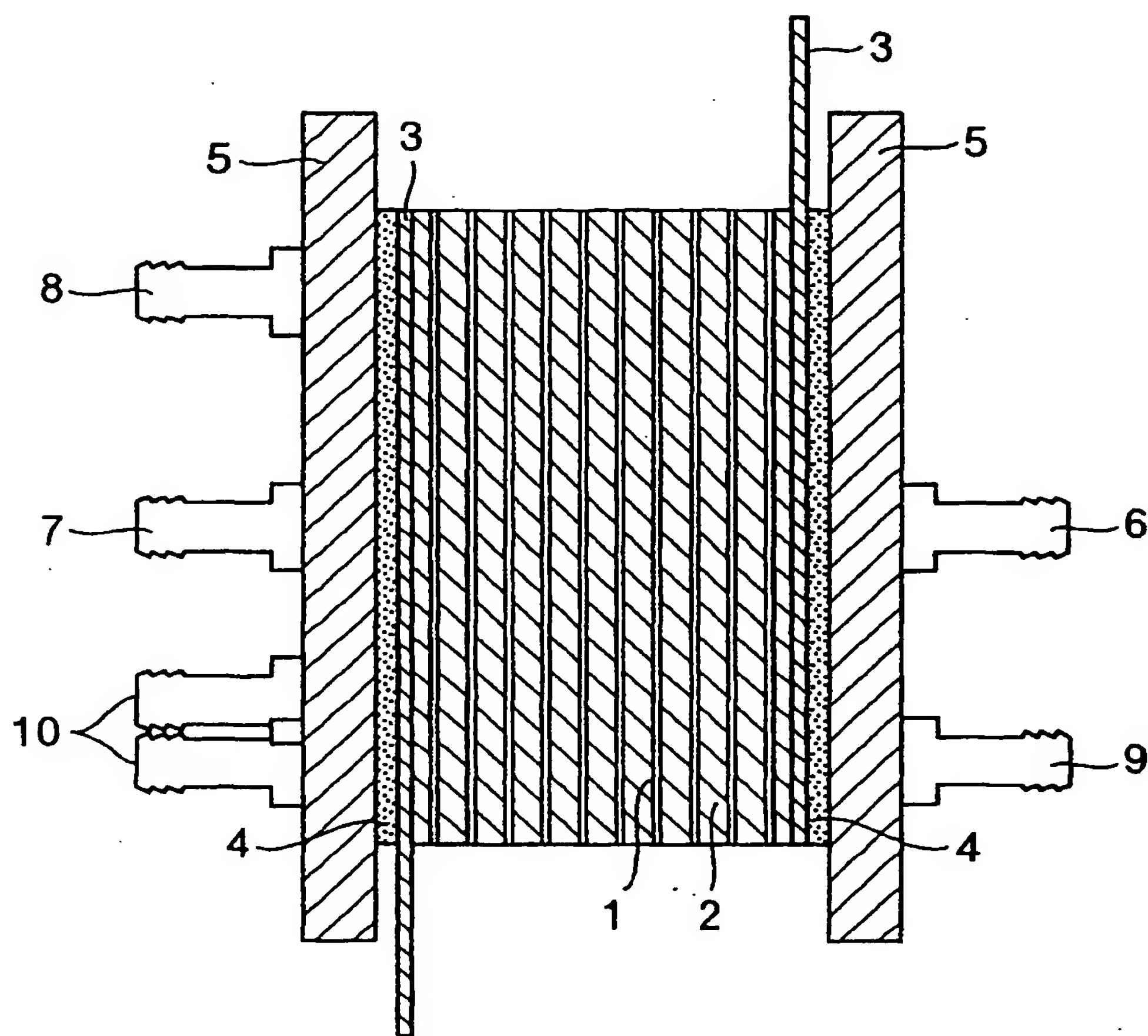


FIG.2





DOCUMENTS CONSIDERED TO BE RELEVANT

| Category | Citation of document with indication, where appropriate, of relevant passages | Relevant to claim | CLASSIFICATION OF THE APPLICATION (Int.Cl.6) |
|---|---|--|--|
| P, X | <p>CHEMICAL ABSTRACTS, vol. 128, no. 18, 4 May 1998 Columbus, Ohio, US; abstract no. 219471, KURITA, TAKESHI ET AL: "Assembled structure of thin fuel cells with good gas-shielding properties" XP002104994 * abstract * & JP 10 055813 A (AISIN SEIKI CO., LTD., JAPAN) -& PATENT ABSTRACTS OF JAPAN vol. 98, no. 6, 30 April 1998 & JP 10 055813 A (AISIN SEIKI CO LTD), 24 February 1998 * abstract * -& DATABASE WPI Derwent Publications Ltd., London, GB; AN 98-203398 XP002104995 A * abstract *</p> <p>—</p> <p>PATENT ABSTRACTS OF JAPAN vol. 018, no. 355 (E-1573), 5 July 1994 -& JP 06 096783 A (MATSUSHITA ELECTRIC IND CO LTD), 8 April 1994 * abstract * -& CHEMICAL ABSTRACTS, vol. 121, no. 4, 25 July 1994 Columbus, Ohio, US; abstract no. 39212, XP002104934 * abstract *</p> <p>—</p> <p>—</p> | 1-3 | |
| A | | 1 | TECHNICAL FIELDS SEARCHED (Int.Cl.6) |
| The present search report has been drawn up for all claims | | | |
| Place of search | Date of completion of the search | Examiner | |
| THE HAGUE | 4 June 1999 | D'hondt, J | |
| CATEGORY OF CITED DOCUMENTS | | T : theory or principle underlying the invention E : earlier patent document, but published on, or after the filing date D : document cited in the application L : document cited for other reasons & : member of the same patent family, corresponding document | |
| X : particularly relevant if taken alone Y : particularly relevant if combined with another document of the same category A : technological background O : non-written disclosure P : intermediate document | | | |



| DOCUMENTS CONSIDERED TO BE RELEVANT | | Relevant to claim | CLASSIFICATION OF THE APPLICATION (Int.Cl.6) |
|---|---|--|--|
| Category | Citation of document with indication, where appropriate, of relevant passages | | |
| A | PATENT ABSTRACTS OF JAPAN vol. 009, no. 251 (E-348), 8 October 1985 & JP 60 101874 A (FUJI DENKI SOUGOU KENKYUSHO:KK), 5 June 1985 * abstract * --- PATENT ABSTRACTS OF JAPAN vol. 008, no. 135 (E-252), 22 June 1984 & JP 59 046767 A (TOKYO SHIBAURA DENKI KK), 16 March 1984 * abstract * --- PATENT ABSTRACTS OF JAPAN vol. 017, no. 450 (E-1416), 18 August 1993 & JP 05 101837 A (MITSUBISHI HEAVY IND LTD), 23 April 1993 * abstract * --- PATENT ABSTRACTS OF JAPAN vol. 007, no. 222 (E-201), 4 October 1983 & JP 58 112269 A (TOKYO SHIBAURA DENKI KK), 4 July 1983 * abstract * --- FR 2 719 946 A (UNITED TECHNOLOGIES CORP) 17 November 1995 * page 6, line 12 - line 35 * * line 30 - line 35; figure 1 * | 1, 3 | |
| A | | 1 | |
| A | | 1 | |
| A | | 3 | |
| A | | 3 | TECHNICAL FIELDS SEARCHED (Int.Cl.6) |
| A | | 3 | |
| The present search report has been drawn up for all claims | | | |
| Place of search | Date of completion of the search | Examiner | |
| THE HAGUE | 4 June 1999 | D'hondt, J | |
| CATEGORY OF CITED DOCUMENTS | | T : theory or principle underlying the invention E : earlier patent document, but published on, or after the filing date D : document cited in the application L : document cited for other reasons & : member of the same patent family, corresponding document | |
| X : particularly relevant if taken alone Y : particularly relevant if combined with another document of the same category A : technological background O : non-written disclosure P : intermediate document | | | |

ANNEX TO THE EUROPEAN SEARCH REPORT
ON EUROPEAN PATENT APPLICATION NO.

EP 99 10 2024

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

04-06-1999

| Patent document cited in search report | Publication date | Patent family member(s) | Publication date |
|--|------------------|-------------------------|------------------|
| FR 2719946 A | 17-11-1995 | NONE | |

EPO FORM P0459

For more details about this annex : see Official Journal of the European Patent Office, No. 12/82